By how much and how quickly does reduction in serum cholesterol concentration lower risk of ischaemic heart disease?

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Abstract

Objective—To estimate by how much and how quickly a given reduction in serum cholesterol concentration will reduce the risk of ischaemic heart disease.

Design—Data on the incidence of ischaemic heart disease and serum cholesterol concentration were analysed from 10 prospective (cohort) studies, three international studies in different communities, and 28 randomised controlled trials (with mortality data analysed according to allocated treatment to ensure the avoidance of bias).

Main outcome measure—Decrease in incidence of ischaemic heart disease or mortality for a 0.6 mmol/l (about 10%) decrease in serum cholesterol concentration.

Results-For men results from the cohort studies showed that a decrease of serum cholesterol concentration of 0.6 mmol/l (about 10%) was associated with a decrease in incidence of ischaemic heart disease of 54% at age 40 years, 39% at age 50, 27% at 60, 20% at 70, and 19% at 80. The combined estimate from the three international studies (for ages 55-64 years) was 38% (95% confidence interval 33% to 42%), somewhat greater than the cohort study estimate of 27%. The reductions in incidence of ischaemic heart disease in the randomised trials (for ages 55-64 years) were 7% (0 to 14%) in the first two years, 22% (15% to 28%) from 2·1-5 years, and 25% (15% to 35%) after five years, the last estimate being close to the estimate of 27% for the long term reduction from the cohort studies. The data for women are limited but indicate a similar effect.

Conclusions—The results from the cohort studies, international comparisons, and clinical trials are remarkably consistent. The cohort studies, based on half a million men and 18 000 ischaemic heart disease events, estimate that a long term reduction in serum cholesterol concentration of 0.6 mmol/l (10%), which can be achieved by moderate dietary change, lowers the risk of ischaemic heart disease by 50% at age 40, falling to 20% at age 70. The randomised trials, based on 45 000 men and 4000 ischaemic heart disease events show that the full effect of the reduction in risk is achieved by five years.

Introduction

In this paper we estimate the size of the reduction in risk of ischaemic heart disease produced by a given reduction in serum cholesterol concentration according to age, and the time needed to attain the full reduction in risk. We use data from observational studies1-14 (adjusted for the two sources of underestimation as described in the preceding paper¹) and from randomised trials. 15-46 The observational studies estimate the maximum expected long term decrease in risk of ischaemic heart disease produced by a given reduction in serum cholesterol concentration since the differences in cholesterol between individual people or between communities will have been present for many years before the data were collected. The randomised trials estimate the extent to which this maximum can be achieved in practice and how quickly.

Methods

ESTIMATING LONG TERM ASSOCIATION—OBSERVATIONAL EVIDENCE

The prospective observational studies of serum cholesterol concentration and ischaemic heart disease fall into two categories. Cohort studies examine the association between ischaemic heart disease and serum cholesterol concentration within cohorts (or groups) of people, generally by comparing subgroups within the cohort defined by ranking individual measurements of cholesterol concentration. International studies compare the incidence of ischaemic heart disease and mean serum cholesterol concentration in separate communities; the comparison is between predefined groups and entails no sorting of individual people by serum cholesterol concentration.

Cohort studies (grouping people by initial cholesterol measurements and comparing rates of ischaemic heart disease)

There are at least 60 cohort studies of serum cholesterol concentration and ischaemic heart disease. many with fewer than 100 ischaemic heart disease events. We confined our attention to the 10 largest published studies, each of which had recorded more than 350 such events (deaths and, in three studies, nonfatal infarcts) in men.1-10 Together they recruited 494 804 men and recorded 18 811 events. The data in each of the studies were published as the incidence of or mortality from ischaemic heart disease in fifths (quintile groups) of the ranked distribution of serum cholesterol concentrations. For each study we regressed the age adjusted rate of ischaemic heart disease (in logarithms) on average cholesterol concentration across the groups. The analysis weighted the rates by the number of events but was not influenced by smoking and blood pressure, which show little association with cholesterol.1245

To estimate the long term effect of a decrease in concentration of low density lipoprotein cholesterol on the risk of ischaemic heart disease from the cohort studies we adjusted for the two sources of underestimation by increasing the slope of the regression line of risk (in logarithms) on total cholesterol concentration by 61%—the estimated correction factor from the BUPA study.1 The two components of this estimate are both generalisable. Similar estimates for regression dilution bias, the larger component, have been obtained from several studies.1 The surrogate dilution effect, the smaller component, will be similar in the 10 studies because total cholesterol values were similar and the concentration of non-low density lipoprotein cholesterol varies little between populations,11 so the proportion of low density lipoprotein cholesterol to total cholesterol would have been similar.

International studies (comparing rates of ischaemic heart disease between communities with varying mean cholesterol concentrations)

The seven countries study measured serum cholesterol concentrations and recorded mortality from ischaemic heart disease prospectively among men living in 16 communities in seven countries.^{12 13} The Ni-Hon-San study similarly compared three commu-

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nities of Japanese men living in Japan, Honolulu, and San Francisco.¹⁴ A third study compared age specific mortality from ischaemic heart disease between 17 countries for which the national average serum cholesterol concentration could be determined from published surveys." For each study we regressed the incidence of or mortality from ischaemic heart disease (in logarithms) in the constituent communities on the cholesterol concentration associated with mean risk in each community. This value is greater than the mean cholesterol concentration (because the association with ischaemic heart disease is log-linear); we estimated it as the mean cholesterol concentration plus a factor, $\frac{1}{2}\sigma^2 b$, calculated from the standard deviation of the concentration in the community, σ , and the regression coefficient, b. The analysis weighted the age adjusted rates for ischaemic heart disease by the number of ischaemic heart disease events and used multiple regression to allow for the confounding effects of smoking and blood pressure. The international studies are not influenced by the underestimation affecting cohort studies. Regression dilution bias is avoided because cholesterol concentrations were not used to divide people into groups, and the differences in mean total cholesterol concentration between communities reflect similar differences in mean low density lipoprotein cholesterol concentration (as the residual nonlow density lipoprotein cholesterol concentration is similar across different communities with widely varying total serum cholesterol concentrations¹¹).

ESTIMATING HOW QUICKLY MAXIMUM REDUCTION IN RISK OF ISCHAEMIC HEART DISEASE CAN BE ATTAINED—RANDOMISED CONTROLLED TRIALS

We identified randomised controlled trials of reduction in cholesterol concentration (by drugs, diet, or ileal bypass surgery) and ischaemic heart disease events (deaths and non-fatal infarcts). We included 28 published trials that recorded at least one death and documented a reduction in serum cholesterol concentration of at least 1%. Together they recruited 46 254 men and recorded 4241 events. Most trials measured only total cholesterol concentration, but the interventions produced similar absolute reductions in concentrations of total and low density lipoprotein cholesterol with little change in non-low density lipoprotein cholesterol. 19 29 31 32 43 46-49 Diagnostic criteria

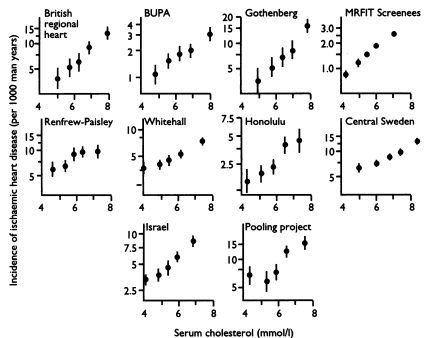


FIG 1—Incidence of ischaemic heart disease, age adjusted with 95% confidence intervals, according to fifths of distribution of serum cholesterol concentration in 10 cohort studies

for death from ischaemic heart disease and non-fatal myocardial infarction were similar in the different trials, and diagnoses were made without knowledge of treatment allocation. Data on mortality were analysed on the basis of allocated treatment, whether or not treatment was received (intention to treat), thereby avoiding bias. With unpublished data on mortality provided by the coordinators of nine of the trials, vital status (alive or dead) at the end of the trial was known for over 99% of all the subjects randomly allocated to interventions in the 28 trials. Data on non-fatal myocardial infarcts were generally not available for subjects who were lost to follow up. Two trials were extended by inviting subjects to continue beyond the planned finishing date1629 but since many declined the random allocation was lost, so we did not include the additional events.

To estimate how quickly the risk of ischaemic heart disease falls after reduction in cholesterol concentration we determined the numbers of ischaemic heart disease events in the trials in three time periods, \leq 2, 2·1-5, and 5·1-12 years after entry. Trial coordinators kindly provided unpublished data for eight trials, and numbers of events were estimated by using published survival curves for two trials.3841 The average reduction in ischaemic heart disease over each time period was estimated by a logistic regression analysis that combined the odds ratios from each trial to obtain a summary relative odds estimate (close to the relative risk as the event rates were low). Each trial was weighted by the mean difference in total cholesterol concentration between treated and control groups. We performed analyses on subgroups of drug and dietary trials and trials of men with and without ischaemic heart disease (angina or previous non-fatal infarction) on entry; authors supplied separate data for two trials that recruited men with and without ischaemic heart disease.34 35

The results of both the observational studies and the trials were expressed as the decrease in the risk of ischaemic heart disease associated with a decrease in serum cholesterol concentration of 0.6 mmol/l; 0.6 mmol/l is about 10% of the average value in Western countries, was the approximate mean reduction in cholesterol concentration attained in the trials, and is a reduction attainable by a moderate reduction in dietary fat.^{47 48}

Results

OBSERVATIONAL STUDIES—MEN Cohort studies

Figure 1 shows the incidence (or mortality) of ischaemic heart disease in the fifths of the ranked cholesterol distribution plotted for each of the 10 cohort studies. The log-linear model for the relation between risk of ischaemic heart disease and serum cholesterol concentration fitted the data well, and in the large study of men screened for inclusion in the multiple risk factor intervention trial (MRFIT screenees) the association was almost perfectly described by a straight line (r=0.997). The magnitude of such a log-linear association can be expressed simply: a constant absolute difference in serum cholesterol concentration, say 0.6 mmol/l, from any point on the cholesterol distribution is associated with a constant percentage difference in the incidence of ischaemic heart disease.

Table I shows this result for each cohort study, unadjusted and adjusted for the two sources of underestimation. In table II the cohorts are subdivided when possible according to age at entry and the results are grouped into 10 year age bands according to the mean age of experiencing ischaemic heart disease events during follow up. The association between ischaemic

					Estimated % decrease in ischaemic heart disease per 0·6 mmol/l decrease in serum cholesterol			
Study	Period of recruitment	Average follow up period (years)	No of subjects	No of deaths from ischaemic heart disease	Unadjusted	Adjusted for underestimation (95% confidence interval)		
	Men							
British regional heart ²	1978-80	8	7 735	438*	22	33 (27 to 39)		
BUPA ¹ (England)	1975-82	13	21 515	538	17	27 (21 to 33)		
Gothenberg (Sweden)	1974-7	7	6 897	360*	26	38 (31 to 44)		
MRFIT screenees (United States)	1973-5	12	361 662	6327	20	30 (28 to 32)		
Renfrew-Paisley' (Scotland)	1972-6	15	7 000	878	11	17 (10 to 22)		
Whitehall (England)	1967-9	18	17718	1542	12	18 (15 to 22)		
Honolulu ⁷	1965-8	19	7 961	371	22	33 (25 to 40)		
Central Sweden ⁸	1963-5	21	46 140	6626	12	19 (17 to 21)		
Israel ⁹	1963	23	9 902	1084	20	30 (25 to 34)		
Pooling project10 (United States)†	1950-60	9	8 274	647*	17	26 (21 to 31)		
	Wome	n						
Renfrew-Paisley ⁵	1972-6	15	8 262	490	12	‡		
Central Sweden ⁸	1963-5	21	46 570	3607	6	‡		

^{*}Deaths and non-fatal infarcts

TABLE II—Estimates (adjusted for underestimation) from 10 cohort studies of percentage decrease in risk of ischaemic heart disease in men per 0.6 mmol/l decrease in serum cholesterol concentration, according to age at death

Study	Estimated % reduction in risk of ischaemic heart disease (95% confidence interval) for age at death (years)								
	35-44	45-54	55-64	65-74	75-84				
British regional heart			33 (27 to 39)						
BUPA		44 (31 to 55)	26 (16 to 36)	19 (10 to 28)					
Gothenberg			38 (31 to 44)						
MRFIT screenees	54 (45 to 62)	38 (33 to 42)	27 (25 to 29)						
Renfrew-Paisley*			17 (5 to 28)	17 (8 to 25)					
Whitehall*		28 (10 to 43)	21 (16 to 26)	16 (11 to 20)					
Honolulu		, ,	• •	33 (25 to 40)					
Central Sweden				, ,	19 (17 to 21				
Israel*			32 (26 to 37)	22 (13 to 29)	•				
Pooling project		40 (30 to 49)	23 (16 to 30)	13 (0 to 25)					
Summary estimate	54	39	27	20	19				

^{*}Unpublished data supplied by authors.

TABLE III—Estimates from international studies of the percentage decrease in incidence of or mortality from ischaemic heart disease per 0.6 mmol/l decrease in serum cholesterol concentration

		No of deaths	Estimated % decrease in ischaemic hear disease per 0·6 mmol/l serum cholestero decrease (95% confidence interval)			
	No of subjects	from ischaemic - heart disease	Men	Women		
Seven countries ¹² 13	11 579	413	36 (23 to 47)	NA		
Ni-Hon-San ¹⁴	11 594	90*	45 (24 to 60)	NA		
International comparison ¹¹	NA	NA	37 (31 to 43)	31 (20 to 40)		
All studies			38 (33 to 42)	31 (20 to 40)		

^{*}Deaths and non-fatal infarcts.

heart disease and serum cholesterol concentration decreases with age. A weighted quadratic regression fitted this relation with age well and yielded estimates that a decrease in cholesterol concentration of 0.6 mmol/l (10%) was associated with a decrease in risk of ischaemic heart disease by 54% at age 40, 39% at age 50, 27% at age 60, 20% at age 70, and 19% at age 80 (table II). A simple combination of the results of the studies in each 10 year age group, weighted inversely by variance, gave almost identical results.

International studies

The three international studies showed similar associations between serum cholesterol concentration and ischaemic heart disease (fig 2, table III). On average a difference in cholesterol concentration of 0.6 mmol/l was associated with a difference in mortality from ischaemic heart disease of 38% (95% confidence interval 33% to 42%) in men. The mean age at death was in the range 55-64 years for all three studies. Differences in serum cholesterol concentration explained over 80% of the international variation in mortality from ischaemic heart disease ($r^2 > 0.80$).

RANDOMISED CONTROLLED TRIALS-MEN

Table IV displays the data from each trial. There was a dose response association, with trials that achieved a greater reduction in serum cholesterol concentration on average showing a greater reduction in ischaemic heart disease (P < 0.001). Table V shows the summary estimates from all the trials combined for a 0.6 mmol/l reduction in cholesterol concentration according to duration of reduction. The reduction in ischaemic heart disease increased with increasing duration of the reduction in cholesterol concentration as follows (for all the trials): within 2 years by 7% (0 to 14%, P=0.06), from 2.1-5 years by 22% (15% to 28%, P < 0.001) and from 5.1-12 years by 25% (15% to 35%, P<0.001), all per 0.6 mmol/l reduction in concentration. There was no significant heterogeneity between the estimates from the different trials within each category of duration (P=0.11). Summary estimates by duration were similar for subgroup analyses of data from the drug and dietary trials and for trials in men with and without known ischaemic heart disease on entry (table V). The overall average reduction was 18% (13% to 22%, P<0.001). The mean range of age at death was 55-64 years in almost all the trials.

In all the trials combined there were highly significant reductions in both fatal ischaemic heart disease and non-fatal myocardial infarction of 10% (3% to 16%, P=0.004) and 21% (15% to 27%, P<0.001) respectively, per 0.6 mmol/l reduction in cholesterol concentration (separate data classified by duration of treatment were not available). The difference between fatal and non-fatal events was significant (P=0.01), but the inference that the risk of non-fatal infarction is reversed more rapidly than that of risk of death from ischaemic heart disease is not justified because the comparison is subject to bias. The analysis of fatal

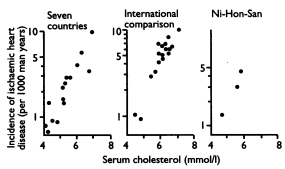


FIG 2—Incidence of ischaemic heart disease according to mean serum cholesterol concentration of different communities in three international studies. (In the seven countries study one community in which only one man died is omitted)

[†]Framingham and four other studies

[‡]Appropriate estimates to adjust for underestimation in women are unavailable.

NA=not applicable.

TABLE IV—Randomised controlled trials of reduction in serum cholesterol concentration. Numbers of men with ischaemic heart disease events (deaths or non-fatal infarcts) by time period

		Mean			No of men	with ischaemi	c heart diseas	e event by time	period sinc	e entr	y to trial
		reduction in serum	No	of men	≤2	Years	2·1-	5 Years	5.	l-12 Y	ears
Trial	Method of cholesterol reduction	cholesterol — concentration (mmol/l)*	Treated	Controls	Treated	Controls	Treated	Controls	Treated		Controls
			Drug	trials							
Men without known ischaemic heart disease:			•								
Helsinki ¹⁵	Gemfibrozil	0.7	2051	2030	27	28	28	53	1		3
World Health Organisation16 17	Clofibrate	0.6	5331	5296	52	56	75	91	46		63
Beggis	Clofibrate	0.7	76	79	4	8	3	11		NA	
Lipid Research Clinic 19†	Cholestyramine	0.7	1906	1900	46	51	51	66	58		70
Men with ischaemic heart disease:			•								
Helsinki ²⁰ †	Gemfibrozil	0.8	311	317	10	14	25	10		NA	
Newcastle ²¹	Clofibrate	0.6	192	208	‡		49	73		NA	
Scottish ²²	Clofibrate	0.6	288	305	÷		51	64		NA	
Coctain	Clofibrate	0.4	1103)	• • • • • • • • • • • • • • • • • • • •	135)		128)		46)		
Coronary drug project23 24†	Cionorato	• •	1100}	2789	}	359	{	372		108	
Coronary urug project	Niacin	0.7	1119	2.07	129	337	126	3.2	32		
Veterans Administration drug-lipid25 26+	Niacin	0.6	145	284	28	48	14	21	22)	NA	
veterans Administration drug-npid	Macm	00	143	201	(years						
Stockholm ²⁷ 28§	Clofibrate, niacin	0.8	279	276	37	41	36	60		NA	
CLAS"	Colestipol, niacin	1.3	94	94	i	5		IA O		NA	
	Colestipol	0.6	23	29	i	ó		IA		NA	
Gross ³⁰ §		0.9	23 57	59 59	3	3	2	7	0	INA	1
NHLBI ¹¹ †	Cholestyramine	1·5	94	59 52	2	0		JA '	U	NA	1
FATS ³²	Three drugs	0.7	79	78	3			IA		NA	
Sahni ³³	Lovastatin	0.7	19	78	3	4	r	iA.		INA	
Men with or without ischaemic heart disease			540	-46		25	_	-		27.4	
Upjohn*†	Colestipol	0.5	548	546	13	25	6	5		NA	
EXCEL*H	Lovastatin	1.1	6582	1663	62	20		ĮA.		NA	
McCaughan ³⁶	Probucol	0.7	88	30	2	2	N	ΙA		NA	
			Dietary	y trials							
Men without known ischaemic heart disease:		0.7	0107	0106			1.4	17		NA	
Minnesota ³⁷		0.7	2197	2196	55	57	14	17		NA	16
Los Angeles ³⁶		0.9	424	422	18	22	23	27	11		16
Men with ischaemic heart disease:	High polyunsaturated								_		_
Medical Research Council**†	low saturated fat diet	1.0	199	194	23	29	22	20	0		3
Oslo ⁴⁰		1.1	229	229	28	34	33	47		NA	
Sydney*'†		0.3	221	237	25	15	12	9		NA	
St Mary's 2 §	Corn oil	0∙6	28	52	8	11		IA.		NA	
STARS*†¶		1.1	60	30	1	4	2	1		NA	
DART#†¶	Low fat diet	0.3	1018	1015	132	144		IA		NA	
London Hospitals 15†		0.6	130	134	39	35	8	15		NA	
			Surger	y trial							
Men with ischaemic heart disease:					24			40			
POSCH*†§	Ileal bypass	1.5	421	417	36	31	15	42	30		53

NA=not applicable

NHLBI=National Heart, Lung and Blood Institute, CLAS=cholesterol-lowering atherosclerosis study, FATS=familial atherosclerosis treatment study, EXCEL=expanded clinical evaluation of lovestatin, STARS=St Thomas's atherosclerosis regression study, DART=diet and reinfarction trial, POSCH=program on surgical control of the hyperlipidemias.

events must have diluted the true magnitude of the effect by including people who did not adhere to their allocated treatment (intention to treat analysis). The analysis of non-fatal events, on the other hand, will have overestimated the true effect through not including such non-compliers. They are typically high risk patients, and since more treated than control patients did not comply, the exclusion of noncompliers will leave the remaining treated patients at lower risk as a group than the remaining control patients. The true effect of reduction of cholesterol concentration is likely to lie between the two estimates for fatal and non-fatal events and is reasonably estimated by our main analysis of a combination of fatal and non-fatal events. In observational studies the strength of the association with serum cholesterol concentration was similar for fatal and non-fatal events.2310

We excluded from our main analysis four trials that used oestrogen and two trials that used thyroxine^{24-26 50-54} to lower serum cholesterol concentration because these two agents have major effects on the risk of ischaemic heart disease that are not mediated through reduction in concentration. Oestrogen produced only a small reduction in serum cholesterol (mean 0·15 mmol/l) and has complex effects on mortality from ischaemic heart disease, reducing it in lower doses but increasing it at higher doses.^{51 55} Thyroxine simulates thyrotoxicosis and so can precipitate myocardial infarction and death from ischaemic heart disease.⁵⁶ Inclusion of these six

trials in the overall analysis changed the estimate of the reduction in ischaemic heart disease per 0.6 mmol/l reduction in cholesterol concentration from 18% (P<0.001) to 14% (P<0.001) but introduced a high degree of heterogeneity between trials (P<0.001), indicating statistical incompatibility between the results of trials that use these two agents and trials that used other regimens to lower cholesterol concentration. Inclusion of four multiple risk factor intervention trials $^{37-60}$ changed the overall result from 18% to 19%.

RESULTS FOR WOMEN

The international estimate was similar for women and men (table III). Three randomised trials with separate data on women showed a significant reduction in ischaemic heart disease similar in size to that observed in men in the same trials.21 22 46 A fourth trial suggested no effect in women, but most of the ischaemic heart disease events occurred within the first few months after reduction of cholesterol concentration.37 Some cohort studies produced smaller estimates of the cholesterol-ischaemic heart disease association in women than in men recruited in the same study,86162 but this can be explained by greater regression dilution bias in women. Serum cholesterol concentration increases steadily in women between the ages of about 35 and 60 years but remains fairly constant in men.11 63 The increase in women would be expected to affect different women to a different extent, so the rank position of individual women in the distribution of

^{*}Difference in mean cholesterol concentration between treated and control subjects throughout trial.

[†]Unpublished data supplied by authors

 $[\]ddagger$ Data for years 0-2 not separately available and are combined with years 2·1-5 years. $\ifmmode Minority$ of subjects are women, sex specific data being unavailable.

^{||}Authors supplied separate data

Some treated patients also took drug cholestyramine.

cholesterol concentrations would change more than in men during the follow up period of the study and regression dilution bias would be greater than in men. Studies with shorter follow up, in which this effect should be less pronounced, recorded similar associations in women and men.⁶¹

Discussion

Our results show a striking consistency between the different categories of data. The cohort studies show the effect of age—a decrease in low density lipoprotein (and total) cholesterol concentration of 0.6 mmol/l in men is associated with a decrease in the risk of ischaemic heart disease of about 50% at the age of 40, 40% at 50, 30% at 60, and 20% at 70 and over (table II). The results of the international studies and randomised trials apply to the age group 55-64 years, and for this age group the estimates from the three types of study can be compared. The cohort study estimate of 27% is lower than the estimate of 38% from the international studies, but the variation in serum cholesterol concentration between countries reflects differences in total dietary fat, reduction of which also favourably alters factor VII activity and coagulation and hence risk of myocardial infarction.64 (Between individual people (the cohort studies) genetic factors are important determinants of the variation in serum cholesterol.)

The trials show that little reduction in risk of ischaemic heart disease occurs in the first two years after lowering cholesterol, but the summary estimate for the reduction in incidence of ischaemic heart disease five or more years after reduction of cholesterol concentration (25%) is close to that from the cohort studies (27%), showing that the full reduction in risk of ischaemic heart disease is achieved within five years. This is consistent with radiographic measurements of the regression of coronary artery atheroma in trials on lowering cholesterol concentration. ²⁹ 31 32 49 46 A man of 35 who reduces his serum cholesterol concentration by 0.6 mmol/l would therefore halve his risk of ischaemic heart disease by the age of 40.

The more limited data available for women indicate that a decrease in serum cholesterol concentration is associated with a similar proportionate reduction in risk of ischaemic heart disease as in men.

The observational data show that in Western societies there is no threshold below which a lower serum cholesterol concentration is not associated with a lower risk of ischaemic heart disease. In the two largest cohort studies the 95% confidence limits of the mortality from ischaemic heart disease in the successive fifths of the cholesterol distribution do not overlap (MRFIT screenees and central Sweden, ¹⁸ fig 1), confirming a continuous relation which, as Chinese data have shown, ⁵⁹ extends to cholesterol values below 4 mmol/l. The international studies show that differences in serum cholesterol concentration (and hence in dietary saturated fat) are the most important determinant of the differences in mortality from ischaemic

TABLE V—Randomised controlled trials of percentage reduction in serum cholesterol concentration: summary estimates of reduction in ischaemic heart disease in men per 0.6 mmol/l (10%) reduction in serum cholesterol concentration according to time period since entry to trial

Trial	Time since entry to trial						
	≤2 years	2·1-5 years	5·1-12 years				
All drug trials	10	21	22				
All dietary trials	9	14	37				
Trials of men without known ischaemic heart disease	11	25	24				
Trials of men with ischaemic heart disease	6	20	26				

All trials (95% confidence interval) 7 (0 to 14) 22 (15 to 28) 25 (15 to 35)

Public health implications

- The combined evidence from the 10 largest cohort studies, three international (ecological) studies, and 28 randomised trials shows conclusively that lowering a person's serum cholesterol concentration results in substantial protection from ischaemic heart disease
- The benefits of serum cholesterol reduction are related to age; a 10% reduction in serum cholesterol concentration produces a reduction in ischaemic heart disease of 50% at age 40, 40% at age 50, 30% at age 60, and 20% at age 70
- The benefit can be realised quickly—the greater part after two years and the full benefit after five years
- Lowering serum cholesterol concentrations in a population is critical in reducing mortality from ischaemic heart disease
- Appropriate action is needed, including wider health education, labelling of foods, and policies on food subsidies that are linked to health priorities

heart disease between countries, accounting for over 80% of the total variation.

An individual person may have difficulty in lowering serum cholesterol concentration by more than about 0.3 mmol/l through dietary change41 44 66; the availability of palatable low fat food may be limited when other members of the family or community do not alter their diet. In high risk patients (those with ischaemic heart disease) cholesterol lowering drugs are justified; they can reduce serum cholesterol by 1.2 mmol/l (20%),35 66 while the dietary changes needed to achieve such a reduction are impractical for an individual person (lowering total dietary fat to about 27% of total energy intake or saturated fat to about 8%).47-49 This 20% reduction in cholesterol would reduce mortality from ischaemic heart disease by half in people aged 55-64 years. On a community basis a reduction in serum concentrations of total and low density lipoprotein cholesterol through dietary change of 0.6 mmol/l (about 10% for total and 15% for low density lipoprotein cholesterol), reducing mortality from ischaemic heart disease by 25-30% in people aged 55-64 years, is a realistic target. It would require a reduction in total dietary fat from about 42% to 35% of total energy intake or a reduction in saturated fat from about 20% to 13%.4748 Reduction in concentration of 0.6 mmol/l in entire communities has occurred through dietary change over periods of a few years11 67 68 and was attained in seven of the nine dietary trials listed in table IV.

Action needs to be taken to reach this target. This includes wider public education, labelling of foods sold in supermarkets, and provision of information on the content of restaurant meals. Most important, though more difficult, is to implement national and international policies on food subsidies that are linked to health priorities.

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- 1 Law MR, Wald NJ, Wu T, Hackshaw A, Bailey A. Systematic underestima-tion of association between serum cholesterol concentration and ischaemic heart disease in observational studies: data from the BUPA study. BM7
- 2 Pocock SJ, Shaper AG, Phillips AN. Concentrations of high density lipo-protein cholesterol, triglycerides, and total cholesterol in ischaemic heart disease. BM7 1989;298:998-1002.
- 3 Rosengren A, Welin L, Tsipogianni A, Wilhelmsen L. Impact of cardiovascular risk factors on coronary heart disease and mortality among m diabetic men: a general population study. *BMJ* 1989;299:1127-31.
- 4 Neaton JD, Wentworth D. Serum cholesterol, blood pressure, smoking, and death from coronary heart disease. Arch Intern Med 1992;152:
- 5 Isles CG, Hole DJ, Hawthorne VM, Lever AF. Relation between coronary risk and coronary mortality in women of the Renfrew and Paisley survey: comparison with men. Lancet 1992;339:702-6
- 6 Shipley MJ, Pocock SJ, Marmot MG. Does plasma cholesterol concentration predict mortality from coronary heart disease in elderly people? 18 year follow up in Whitehall study. BMJ 1991;303:89-92.
- Stemmermann GN, Chyou P-H, Kagan A, Nomura AMY, Yano K. Serum cholesterol and mortality among Japanese-American males: the Honolulu (Hawaii) heart program. Arch Intern Med 1991;151:969-72.
- 8 Tornberg SA, Holm L-E, Carstensen JM, Eklund GA. Cancer incidence and cancer mortality in relation to serum cholesterol. J Natl Cancer Inst 1989;81:1917-21.
- 9 Goldbourt U, Yaari S. Cholesterol and coronary heart disease mortality: a 23 year follow-up study of 9,902 men in Israel. Arteriosclerosis 1990;10:512-9.
- 10 The Pooling Project Research Group. Relationship of blood pressure, serum cholesterol, smoking habit, relative weight and ECG abnormalities to incidence of major coronary events: final report of the Pooling Project. *Journal of Chronic Diseases* 1978;31:201-306.
- 11 Law MR, Wald NJ. An ecological study of serum cholesterol and ischaemic
- heart disease between 1950 and 1990. Eur J Clin Nutr (in press).

 12 Keys A. Seven Countries: a multivariate analysis of death and coronary heart
- disease. Cambridge: Harvard University Press, 1980.

 13 Keys A, Aravanis C, Blackburn H, van Buchem F, Buzina R, Djordjevic B, et al. Epidemiological studies related to coronary heart disease: charac teristics of men aged 40-59 in seven countries. Acta Med Scand 1967; suppl 480:1-392.
- 14 Robertson TL, Kato H, Rhoads GG, Kagan A, Marmot M, Syme SL, et al. Epidemiologic studies of coronary heart disease and stroke in Japanese men living in Japan, Hawaii and California. Am J Cardiol 1977;39:239-49
- 15 Frick MH, Elo O, Haapa K, Heinonen OP, Heinsalmi P, Helo P, et al. Helsinki heart study: primary prevention trial with gemfibrozil in middle-aged men with dyslipidemia. N Engl J Med 1987;317:1237-45.
- committee of Principal Investigators. A co-operative trial in the primary prevention of ischaemic heart disease using clofibrate. Br Heart J 1978;40: 1069-118
- 17 Heady JA, Morris IN, Oliver MF, WHO clofibrate/cholesterol trial: clarifica-
- tions (letter). Lancet 1992;340:1405-6.

 18 Begg TB, Rifkind BM. Valutazione della terapia con clofibrate nelle arterioe periferiche. Minerva Med 1971;62:3469-75
- 19 Lipid Research Clinics Program. The Lipid Research Clinics coronary primary evention trial results. I. Reduction in incidence of coronary heart disease. 7AMA 1984:251:351-64.
- 20 Frick MH, Heinonen OP, Huttunen JK, Koskinen P, Mänttäri M, Manninen V. Efficacy of gemfibrozil in dyslipidaemic subjects with suspected heart disease. An ancillary study in the Helsinki Heart Study Frame Population. Ann Med 1993;25:41-5.
- 21 Group of Physicians of the Newcastle upon Tyne Region. Trial of clofibrate in the treatment of ischaemic heart disease. BMJ 1971;iv:767-75.
- 22 Research Committee of the Scottish Society of Physicians. Ischaemic heart disease: a secondary prevention trial using clofibrate. BMJ 1971;iv:775-84.
- 23 Coronary Drug Project Research Group. Clofibrate and niacin in coronary heart disease. JAMA 1975;231:360-81.
- 24 Canner PL, Berge KG, Wenger NK, Stamler J, Friedman L, Prineas RJ, et al. Fifteen year mortality in coronary drug project patients: long-term benefit with niacin. J Am Coll Cardiol 1986;8:1245-55.
 25 Schoch HK. The US Veterans Administration cardiology drug-lipid study: an
- interim report. Adv Exp Med Biol 1968;4:405-20.
- 26 Detre KM, Shaw L. Long-term changes of serum cholesterol with cholesterolaltering drugs in patients with coronary heart disease. Veterans Administration drug-lipid cooperative study. Circulation 1974;50:998-1005.
- 27 Carlson I.A, Rosenhamer G. Reduction of mortality in the Stockholm ischaemic heart disease secondary prevention study by combined treatment with clofibrate and nicotinic acid. Acta Med Scand 1988;223:405-18.
- 28 Rosenhamer G, Carlson LA. Effect of combined clofibrate-nicotinic acid treatment in ischaemic heart disease. *Atherosclerosis* 1980;37:129-38.

 29 Blankenhorn DH, Nessim SA, Johnson RL, Sanmarco ME, Azen SP, Cashin-
- Hemphill L. Beneficial effects of combined colestipol-niacin the coronary atherosclerosis and coronary venous bypass grafts. JAMA 1987; 257:3233-40
- 30 Gross L, Figueredo R. Long-term cholesterol-lowering effect of colestipol resin in humans. 7 Am Geriatr Soc 1973;21:552-6.
- 31 Brensike JF, Levy RI, Kelsey SF, Passamani ER, Richardson JM, Loh IK, et al. Effects of therapy with cholestyramine on progression of coronary arteriosclerosis: results of the NHLBI type II coronary intervention study. Circulation 1984:69:313-24.
- 32 Brown G, Albers JJ, Fisher LD, Schaefer SM, Lin J-T, Kaplan C, et al. Regression of coronary artery disease as a result of intensive lipid-lowering therapy in men with high levels of apolipoprotein B. N Engl J Med 1990;323:1289-98.
- 33 Sahni R, Maniet AR, Voci G, Banka VS. Prevention of restenosis by lovastatin after successful coronary angioplasty. Am Heart J 1991;121:1600-8.
 34 Dorr AE, Gundersen K, Schneider JC, Spencer TW, Martin WB. Colestipol
- hydrochloride in hypercholesterolemic patients—effect on serum cholesterol and mortality. *Journal of Chronic Diseases* 1978;31:5-14.
- 35 Bradford RH, Shear CL, Chremos AN, Dujovne C, Downton M. Franklin FA, et al. Expanded clinical evaluation of lovastatin (EXCEL) study results.

- I. Efficacy in modifying plasma lipoproteins and adverse event profile in 8,245 patients with moderate hypercholesterolemia. Arch Intern Med
- 36 McCaughan D. The long-term effects of probucol on serum lipid levels. Arch Intern Med 1981;141:1428-32.
- 37 Frantz ID, Dawson EA, Ashman PL, Gatewood LC, Bartsch GE, Kuba K, al. Test of effect of lipid lowering by diet on cardiovascular risk. The Minnesota coronary survey. Arteriosclerosis 1989:9:129-35.
- 38 Dayton S, Pearce ML, Hashimoto S, Dixon WJ, Tomiyasu U. A controlled clinical trial of a diet high in unsaturated fat. Circulation 1969;39-40(suppl II):II1-63.
- 39 Research Committee to the Medical Research Council. Controlled trial of sovabean oil in myocardial infarction. Lancet 1968ii:693-700.
- 40 Leren P. The effect of plasma cholesterol lowering diet in male survivors of myocardial infarction. Acta Med Scand 1966;suppl 466:1-92.
 41 Woodhill JM, Palmer AJ, Leelarthaepin B, McGilchrist C, Blacket RB. Low
- fat, low cholesterol diet in secondary prevention of coronary heart disease. Adv Exp Med Biol 1978;109:317-30.
- 42 Rose GA, Thomson WB, Williams RT. Corn oil in treatment of ischaemic heart disease. BM7 1965;i:1531-3.
- 43 Watts GF, Lewis B, Brunt JNH, Lewis ES, Coltart DJ, Smith LDR, et al. Effects on coronary artery disease of lipid-lowering diet, or diet plus cholestyramine, in the St Thomas' Atherosclerosis Regression Study (STARS). Lancet 1992;339:563-9.
- urr ML, Fehily AM, Gilbert JF, Rogers S, Holliday RM, Sweetnam PM, et al. Effects of changes in fat, fish and fibre intakes on death and myocardial reinfarction: diet and reinfarction trial (DART). Lancet 1989;ii:757-61
- 45 Research Committee (from three London hospitals). Low fat diet in myocardial infarction. Lancet 1965;ii:501-4.
- 46 Buchwald H, Varco RL, Matts JP, Long JM, Fitch LL, Campell GS, et al. Effect of partial ileal bypass surgery on mortality and morbidit coronary heart disease in patients with hypercholesterolemia. N Engl J Med 1990;323:946-55.
- 47 Grundy SM, Nix D, Whelan MF, Franklin L. Comparison of three cholesterol-lowering diets in normolipidemic men. JAMA 1986;256: 2351-5
- 48 Watts GF, Ahmed W, Quiney J, Houlston R, Jackson P, Iles C, et al. Effective
- lipid lowering diets including lean meat. BMJ 1988;296:235-7.
 49 Lewis B, Hammett F, Katan M, Kay RM, Merkx I, Nobels A, et al. Towards an improved lipid-lowering diet: additive effects of changes in nutrient intake. Lancet 1981;ii:1310-3.
- 50 Marmorston I, Moore FI, Hopkins CE, Kuzma OT, Weiner I. Clinical studies of long-term estrogen therapy in men with myocardial infarction. *Proc Soc Exp Biol Med* 1962;110:400-8.
- 51 Stamler J, Pick R, Katz LN, Pick A, Kaplan BM, Berkson DM, et al, Effectiveness of estrogens for therapy of myocardial infarction in middle-age men. JAMA 1963;183:632-8.
- 52 The Coronary Drug Project Research Group. The Coronary Drug Project initial findings leading to modifications of its research protocol. JAMA 1970;214:1303-13.
- 53 The Coronary Drug Project Research Group. The Coronary Drug Project: findings leading to discontinuation of the 2-5 mg/day estrogen group. JAMA 1973-226-652-7
- 54 The Coronary Drug Project Research Group. Findings leading to further modifications of its protocol with respect to dextrothyroxine. JAMA 1972;220:996-1008.
- 55 Henriksson P, Edhag O. Orchidectomy versus oestrogen for prostatic cancer: cardiovascular effects. BMJ 1986;293:413-5.
- 56 Reynolds JEF, ed. Martindale: the Extra Pharmacopoeia. London: Pharmaceutical Press, 1989.
- 57 Multiple Risk Factor Intervention Trial Research Group. Multiple risk factor intervention trial. Risk factor changes and mortality results. JAMA 1982:248:1465-77.
- 58 Hjermann I, Holme I, Velve Byre K, Leren P. Effect of diet and smoking intervention on the incidence of coronary heart disease. Lancet 1981;ii:
- 59 Miettinen TA, Huttunen JK, Naukkarinen V, Strandberg T, Mattila S, Kumlin T, et al. Multifactorial primary prevention of cardiovascular diseases in middle-aged men: risk factor changes, incidence, and mortality. JAMA 1985;254:2097-102.
- 60 Singh RB, Rastogi SS, Verma R, Laxmi B, Singh R, Ghosh S, et al. Randomised controlled trial of cardioprotective diet in patients with recent acute myocardial infarction: results of one year follow up. BMJ 1992;304:
- 61 Crouse JR. Gender, lipoproteins, diet, and cardiovascular risk. Lancet 1989;i:
- 62 Manolio TA, Pearson TA, Wenger NK, Barrett-Connor E, Payne GH, Harlan WR. Cholesterol and heart disease in older persons and women: review of an NHLBI Workshop. Annals of Epidemiology 1992;2:161-76.
 63 The Lipid Research Clinics Program Epidemiology Committee. Plasma lipid
- distributions in selected North American populations: the Lipid Research Clinics program prevalence study. Circulation 1979;60:427-39.

 64 Miller GJ, Cruickshank JK, Ellis LJ, Thompson RL, Wilkes HC, Stirling Y, et al. Fat consumption and factor VII coagulant activity in middle-aged men. An association between a dietary and thrombogenic coronary risk factor. Atherosclerosis 1989;78:19-24
- 65 Chen Z, Peto R, Collins R, MacMahon S, Lu J, Li W. Serum cholesterol concentration and coronary heart disease in a population with low cholesterol concentrations. BM7 1991:303:276-82.
- 66 Hunninghake DB, Stein EA, Dujovne CA, Harris WS, Feldman EB, Miller VT, et al. The efficacy of intensive dietary therapy alone or combined with lovastatin in outpatients with hypercholesterolemia. N Engl J Med 1993; 328:1213-9
- Sigfusson N, Sigvaldason H, Steingrimsdottir L, Gudmundsdottir II, Stafansdottir I, Thorsteinsson T, et al. Decline in ischaemic heart disease in Iceland and change in risk factor levels. BMJ 1991;302:1371-5
- 68 Puska P, Salonen JT, Nissinen A, Tuomilehto J, Vartiainen E, Korhonen H, et al. Change in risk factors for coronary heart disease during 10 years of a community intervention programme (north Karelia project). BMJ 1983; 287:1840-4.

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